WEST Search History

09/664 186 A##

DATE: Friday, May 24, 2002

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61	L10
46671	L9
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7	L6
1	L5
7	L4
921	L3
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1963	L1
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Search Results - Record(s) 1 through 7 of 7 returned.

1. <u>6350591</u> . 16 Feb 99; 26 Feb 02. Recombinant DNA and methods for producing thermostable enzymes. Weber; J. Mark, et al. 435/69.1; 435/477 536/23.7. C12N015/74 C12N015/31.			
2. <u>6344327</u> . 12 Apr 00; 05 Feb 02. Methods for isolation of thermophile promoters. Peredultchuk; Mikhail, et al. 435/6; 435/252.3 435/29 435/440 435/471 435/477 435/69.1 536/23.1 536/24.1. C12Q001/68 C12Q001/02 C12N001/20 C12N015/00 C07H021/04.			
3. <u>6294358</u> . 07 Sep 99; 25 Sep 01. Thermus promoters for gene expression. Peredultchuk; Mikhail, et al. 435/69.1; 435/252.3 435/320.1 435/440 435/477 435/6 536/23.1 536/24.1. C12P021/00 C12N015/00 C12N015/74 C07H021/04.			
4. <u>6207377</u> . 14 Aug 98; 27 Mar 01. Method for construction of <u>thermus</u> -E. coli <u>shuttle</u> vectors and identification of two <u>Thermus</u> plasmid replication origins. Wayne; Jay, et al. 435/6; 435/252.3 435/320.1 435/471 435/91.1 536/23.1 536/24.1. C12Q001/68 C12P019/34 C12N015/74 C12N015/63 C12N001/20.			
5. <u>5872238</u> . 18 Aug 97; 16 Feb 99. Thermophile gene transfer. Weber; J. Mark, et al. 536/23.7;. C12N015/31.			
6. <u>5786174</u> . 28 Jan 97; 28 Jul 98. Thermophile gene transfer. Weber; J. Mark, et al. 435/69.1; 435/463 530/350 536/23.1. C12P021/02 C12N015/63 C07K014/00 C07H021/04.			
7. <u>5120658</u> . 28 Mar 89; 09 Jun 92. Thermostable tryptophan synthetase gene and extremely thermophilic plasmid vector incorporating said gene. Koyama; Yoshinori, et al. 435/320.1; 435/108 435/183 435/252.3 435/69.1 435/71.2 435/91.41 536/23.2. C12N015/70 C12N015/52.			
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1. Document ID: US 6350591 B1

L6: Entry 1 of 7

File: USPT

Feb 26, 2002

US-PAT-NO: 6350591

DOCUMENT-IDENTIFIER: US 6350591 B1

TITLE: Recombinant DNA and methods for producing thermostable enzymes

DATE-ISSUED: February 26, 2002

US-CL-CURRENT: 435/69.1; 435/477, 536/23.7

APPL-NO: 9/ 250585

DATE FILED: February 16, 1999

PARENT-CASE:

This application is a continuation of application Ser. No. 08/912,794, filed, Aug. 18, 1997, now U.S. Pat. No. 5,872,238, which is a continuation of application Ser. No. 08/496,932 filed Jun. 30, 1995, now abandoned, which is a continuation of application Ser. No. 08/265,522, filed Jun. 24, 1994, now abandoned.

IN: Weber; J. Mark, Demirjian; David C., Casadaban; Malcolm J., Vonstein; Veronika, Pagratis; Nikos C.

AB: We have developed a new gene transfer system for extreme thermophiles of the genus Thermus, including

Thermus flavus., using a chromosomal gene, and a thermostable derivative of the kanamycin-resistance gene

(kan.sup.tr2). A plasmid mediated gene-replacement process is used to insert it into the chromosome resulting in the

production of Leu.sup.- Km.sup.r transformants. This system not only allows stable, single-copy gene insertion into

the chromosome of an extreme thermophile, but can be used in the thermo-genetic process described here to generate

thermo-stabilized enzymes and proteins for industrial processes. This host-vector environment makes it possible to

generate further thermo-stabilizing mutations in the kan gene beyond those levels previously reported.

L6: Entry 1 of 7

File: USPT

Feb 26, 2002

DOCUMENT-IDENTIFIER: US 6350591 B1

TITLE: Recombinant DNA and methods for producing thermostable enzymes

Brief Summary Paragraph Right (9):

Koyama et al. (1990) A plasmid vector for an extreme thermophile, Thermus thermophilus, FEMS Microbiology Letters 72:97-102, teach a Thermus-E. coli shuttle vector carrying a tryptophan synthetase gene (trpB). This cryptic plasmid pTT8, was able to transform Thermus thermophilus. The authors point out that a plasmid vector carrying trpBA was not suitable for selection since the cloned DNA fragment recombined with the chromosomal counterpart at high frequency.

Brief Summary Paragraph Right (17):

Lasa et al. (1992a) Development of Thermus-Escherichia Shuttle Vectors and Their Use for Expression of the Clostridium thermocellum celA Gene in Thermus thermophilus, J. of Bacteriology 174:6424-6431, teach the self-selection of undescribed origins of replication from cryptic plasmids from uncharacterized Thermus spp. and Thermus aquaticus are isolated and cloned into E. coli vectors. Plasmids were constructed with these origins, pLU1 to pLU4 from T. aquaticus, and pMY1 to pMY3 from Thermus spp. The plasmids then had a modified form of the cellulase gene (celA) from

Clostridium thermocellum and were expressed in E. coli with the signal peptide from the S-layer gene from T. thermophilus. Transformation back into T. thermophilus allowed for expression at 70.degree. C.

2. Document ID: US 6344327 B1

L6: Entry 2 of 7

File: USPT

Feb 5, 2002

US-PAT-NO: 6344327

DOCUMENT-IDENTIFIER: US 6344327 B1

TITLE: Methods for isolation of thermophile promoters

DATE-ISSUED: February 5, 2002

US-CL-CURRENT: 435/6; 435/252.3, 435/29, 435/440, 435/471, 435/477, 435/69.1, 536/23.1, 536/24.1

APPL-NO: 9/548260 DATE FILED: April 12, 2000

PARENT-CASE:

This application is a divisional application of U.S. patent application Ser. No. 09/390,867, filed Sep. 7, 1999.

IN: Peredultchuk; Mikhail, Vonstein; Veronica, Demirjian; David

AB: The present invention relates to a system for identifying, isolating and utilizing promoter elements

useful for expression of nucleotide sequences and the proteins encoded thereby in a thermophile. In one embodiment,

a recombinant DNA molecule is provided, and comprises a reporter sequence, a putative thermophile promoter, a

selectable marker sequence, and a 3' and a 5' DNA targeting sequence that are together capable of causing

integration of at least a portion of said DNA molecule into the genome of a thermophile. Further, within the

recombinant DNA, the reporter sequence is under the transcriptional control of a promoter which functions in a

thermophile to form a promoter/reporter cassette, the promote/reporter cassette is flanked by said 3' and said 5'

DNA targeting sequences, and the promoter/reporter cassette is positioned in the opposite orientation of the DNA targeting sequences.

L6: Entry 2 of 7

File: USPT

Feb 5, 2002

DOCUMENT-IDENTIFIER: US 6344327 B1
TITLE: Methods for isolation of thermophile promoters

Drawing Description Paragraph Right (3): FIG. 3. Construction of pTG200 and development of promoter test vectors. A) Comparison of terminator sequences from Thermus. The his terminator was used in the construction of pTG200. B) pTG200 consists of an E. coli shuttle vector with the Thermus leuB gene

disrupted by the promoterless kantr2 gene in the opposite direction. A strong .

Thermus transcription terminator is placed ownstream

of the Kantr2 gene to prevent transcription through the gene in the opposite

direction. Promoter-test vectors were constructed by using primers to the two ends of the kan gene with an extended 50-60 bp promoter attached at the 5' end. Leu terminator (SEQ ID NO. 47); his terminator (SEQ ID NO. 48); icd terminator (SEQ ID NO. 49); proC terminator (SEQ ID NO. 50); phe S/T terminator (SEQ ID NO. 51); pol terminator (SEQ ID NO. 52).

Detailed Description Paragraph Right (5):
Using the reagents and techniques described in this application, inducible and constitutive promoters, integrative and plasmid-based vectors, and nucleic acids containing secretion signals may be isolated. The vectors utilized may be any vector suitable to isolation and characterization of a promoter. For instance, the vectors utilized may be plasmid, bacteriphage, virus, phagemid, cointegrate of one or more species, etc. Preferably, the vector is amenable to expression of a nucleotide sequence in a prokaryotic cell such as Thermus or E. coli. It is further preferable that the vectors be capable of functioning in different

Detailed Description Paragraph Right (10):

types of cells (ie, shuttle), such as Thermus or E coli.

Liao, et al. ((1986) Isolation of a thermostable enzyme variant by cloning and selection in a thermophile. Proc. Natl. Acad. Sci.

USA. 83:576-580) first demonstrated in vivo thermostabilization of a gene by using kanamycin nucleotidyl transferase in Bacillus

stearothermophilus where resistance to 63.degree. C. was shown. To improve the genetic thermostabilization approach, a gene

transfer system for Thermus was developed where the upper growth limit was above 80.degree. C. instead of 65.degree. C. as in

Bacillus (described in, for example, U.S. Pat. No. 5,786,174 which is hereby incorporated by reference). These experiments were

intitially conducted using the thermostabilized kan gene, in which the initial Km.sup.r supported growth only to 55.degree. C. in

Thermus and not to 63.degree. C. as reported by Liao, et. al. ((1986) Isolation of a thermostable enzyme variant by cloning and

selection in a thermophile. Proc. Natl. Acad. Sci. USA 83:576-580). The regulated expression system provided herein allows for

fine-tuning of thermostabilization selection experiments so that the temperature range can be regulated and controlled and cutoff temperatures for selection adjusted in subsequent rounds of mutagenesis.

Some important elements of Thermus' genetic background have been previously described. The generation of mutations (Koyama, et al.

(1990) Cloning and sequence analysis of tryptophan synthetase genes of an extreme thermophile, Thermus thermophilus HB27: Plasmid transfer from replica-plated Escherichia coli

recombinant colonies to competent T. thermophilus cells. J. Bacteriol. 172:3490-3495; Koyama, et al. (1990) A plasmid vector for

an extreme thermophile, Thermus thermophilus. FEMS Microbiol. Lett. 72:97-102; Lasa, et al. (1992) Insertional mutagenesis in the

extreme thermophilic eubacteria Thermus thermophilus HB8. Molec. Microbiol. 6:1555-1564), chromosomal integration (Koyama, et al. (1990) Cloning and sequence analysis of tryptophan synthetase genes of an

extreme thermophile, Thermus thermophilus HB27: Plasmid transfer from replica-plated Escherichia coli recombinant colonies to competent T. thermophilus cells. J. Bacteriol. 172:3490-3495; Koyama, et al. (1990) A plasmid vector for an extreme

172.3490-3495; Koyama, et al. (1990) A plasmid vector for an extreme thermophile, Thermus thermophilus. FEMS Microbiol Lett. 72:97-102; Lasa, et al. (1992) Insertional mutagenesis in the extreme

thermophilic eubacteria Thermus thermophilus HB8. Molec. Microbiol. 6:1555-1564), plasmids (Mather, et al. (1990) Plasmid-associated

aggregation in Thermus thermophilus HB8. Plasmid. 24:45-56; Hishinuma, et al. (1978) Isolation of extrachromosomasl deoxyribonucleic Acids from extremely thermophilic bacteria.

deoxyribonucleic Acids from extremely thermophilic bacteria.

Jour. of General Microbiology. 104:193-199.). and phages (Sakaki, et al. (1975) Isolation and Characterization of a Bacteriophage

Infectious to an Extreme Thermophile. Thermus thermophilus HB8. J. Virol. 15:1449-1453) have also been studied. Several successful

attempts to develop cloning systems using plasmids and chromosomal integration systems were demonstrated (Koyama, et al. (1986)

Genetic transformation of the extreme thermophile thermos thermophilus and of other thermos spp. J. Bacteriol. 166:338-340; Lasa, et al. (1992) Development of Thermus-Escherichia Shuttle Vectors and Their

Use for Expression of the Clostridium thermocellum celA Gene in Thermus thermophilus. J. Bacteriol. 174:6424-6431; Mather, et al.

(1992) Development of Plasmid Cloning Vectors for Thermus thermophilus HB8: Expression of a Heterologous, Plasmid-Borne Kanamycin Nucleotidyltransferase Gene. Appl. Environ.

Microbiol 58:421-425.). However, none of these provide the versatility as

those provided herein.

Other Reference Publication (14):

Lasa et al., "Development of Thermus-Escherichia Shuttle Vectors and Their Use for Expression of the Clostridium thermocellum celA Gene in Thermus thermophilus," J. of Bacteriology, 174:6424-6431 (1992a).

3. Document ID: US 6294358 B1

L6: Entry 3 of 7

File: USPT

Sep 25, 2001

US-PAT-NO: 6294358

DOCUMENT-IDENTIFIER: US 6294358 B1

TITLE: Thermus promoters for gene expression

DATE-ISSUED: September 25, 2001

US-CL-CURRENT: 435/69.1; 435/252.3, 435/320.1, 435/440, 435/477, 435/6, 536/23.1, 536/24.1

APPL-NO: 9/ 390867

DATE FILED: September 7, 1999

IN: Peredultchuk; Mikhail, Vonstein; Veronica, Demirjian; David C.

AB: The present invention relates to a system for identifying, isolating and utilizing promoter elements

useful for expression of nucleotide sequences and the proteins encoded thereby in a thermophile. In one embodiment,

a recombinant DNA molecule is provided, and comprises a reporter sequence, a putative thermophile promoter, a

selectable marker sequence, and a 3' and a 5' DNA targeting sequence that are together capable of causing

integration of at least a portion of said DNA molecule into the genome of a thermophile. Further, within the

recombinant DNA, the reporter sequence is under the transcriptional control of a promoter which functions in a

thermophile to form a promoter/reporter cassette, the promote/reporter cassette is flanked by said 3' and said 5'

DNA targeting sequences, and the promoter/reporter cassette is positioned in the opposite orientation of the DNA targeting sequences.

L6: Entry 3 of 7

File: USPT

Sep 25, 2001

DOCUMENT-IDENTIFIER: US 6294358 B1 TITLE: Thermus promoters for gene expression

Drawing Description Paragraph Right (3):

FIG. 3. Construction of pTG200 and development of promoter test vectors.

A) Comparison of terminator sequences from Thermus SEQ ID

NO: 47-SEQ ID NO: 52. The his terminator was used in the construction of pTG200. B) pTG200 consists of an E. coli shuttle vector

with the Thermus leuB gene disrupted by the promoterless kantr2 gene in the opposite direction. A strong Thermus transcription

terminator is placed downstream of the kantr2 gene to prevent transcription through the gene in the opposite direction.

Promoter-test vectors were constructed by using primers to the two ends of the kan gene with an extended 50-60 bp promoter

attached at the 5'end.

Detailed Description Paragraph Right (5):
Using the reagents and techniques described in this application, inducible and constitutive promoters, integrative and plasmid-based vectors, and nucleic acids containing secretion signals may be isolated. The vectors utilized may be any vector suitable to isolation and characterization of a promoter. For instance, the vectors utilized may be plasmid, bacteriophage, virus, phagemid, cointegrate of one or more species, etc. Preferably, the vector is amenable to expression of a nucleotide sequence in a prokaryotic cell such as Thermus or E. coli. It is further preferable that the vectors be capable of functioning in different types of cells (ie, shuttle), such as Thermus or E. coli.

Detailed Description Paragraph Right (10): Liao, et al. ((1986) Isolation of a thermostable enzyme variant by cloning and selection in a thermophile. Proc. Natl. Acad. Sci. USA. 83:576-580) first demonstrated in vivo thermostabilization of a gene by using kanamycin nucleotidyl transferase in Bacillus stearothermophilus where resistance to 63.degree. C. was shown. To improve the genetic thermostabilization approach, a gene transfer system for Thermus was developed where the upper growth limit was above 80.degree. C. instead of 65.degree. C. as in Bacillus (described in, for example, U.S. Pat. No. 5,786,174 which is hereby incorporated by reference). These experiments were intitially conducted using the thermostabilized kan gene, in which the initial Km.sup.r supported growth only to 55.degree. C. in Thermus and not to 63.degree. C. as reported by Liao, et. al. ((1986) Isolation of a thermostable enzyme variant by cloning and selection in a thermophile. Proc. Natl. Acad. Sci. USA. 83:576-580). The regulated expression system provided herein allows for fine-tuning of thermostabilization selection experiments so that the temperature range can be regulated and controlled and cutoff temperatures for selection adjusted in subsequent rounds of mutagenesis. Some important elements of Thermus' genetic background have been previously described. The generation of mutations (Koyama, et al. (1990) Cloning and sequence analysis of tryptophan synthetase genes of an extreme thermophile, Thermus thermophilus HB27: Plasmid transfer from replica-plated Escherichia coli recombinant colonies to competent T. thermophilus cells. J. Bacteriol. 172:3490-3495; Koyama, et al. (1990) A plasmid vector for an extreme thermophile, Thermus thermophilus. FEMS Microbiol. Lett. 72:97-102; Lasa, et al. (1992) Insertional mutagenesis in the extreme thermophilic eubacteria Thermus thermophilus HB8. Molec. Microbiol. 6:1555-1564), chromosomal integration (Koyama, et al. (1990) Cloning and sequence analysis of tryptophan synthetase genes of an extreme thermophile, Thermus thermophilus HB27: Plasmid transfer from replica-plated Escherichia coli recombinant colonies to competent T. thermophilus cells. J. Bacteriol. 172:3490-3495; Koyama, et al. (1990) A plasmid vector for an extreme thermophile, Thermus thermophilus. FEMS Microbiol. Lett. 72:97-102; Lasa, et al. (1992) Insertional mutagenesis in the extreme thermophilic eubacteria Thermus thermophilus HB8. Molec. Microbiol. 6:1555-1564), plasmids (Mather, et al. (1990) Plasmid-associated aggregation in Thermus thermophilus HB8. Plasmid. 24:45-56; Hishinuma, et al. (1978) Isolation of extrachromosomasl deoxyribonucleic Acids from extremely thermophilic bacteria. Jour. of General Microbiology. 104:193-199.), and phages (Sakaki, et al. (1975) Isolation and Characterization of a Bacteriophage Infectious to an Extreme Thermophile, Thermus thermophilus HB8. J. Virol. 15:1449-1453) have also been studied. Several successful attempts to develop cloning systems using plasmids and chromosomal integration systems were demonstrated (Koyama, et al. (1986) Genetic transformation of the extreme thermophile thermos thermophilus and of other thermus spp. J. Bacteriol. 166:338-340; Lasa, et al. (1992) Development of Thermus-Escherichia Shuttle Vectors and Their Use for Expression of the Clostridium thermocellum celA Gene in Thermus thermophilus. J. Bacteriol. 174:6424-6431; Mather, et al. (1992) Development of Plasmid Cloning Vectors for Thermus thermophilus HB8: Expression of a Heterologous, Plasmid-Borne Kanamycin Nucleotidyltransferase Gene. Appl. Environ. Microbiol 58:421-425.). However, none of these provide the versatility as

those provided herein.

4. Document ID: US 6207377 B1

L6: Entry 4 of 7

File: USPT

Mar 27, 2001

US-PAT-NO: 6207377 DOCUMENT-IDENTIFIER: US 6207377 B1

TITLE: Method for construction of thermus-E. coli shuttle vectors and identification of two Thermus plasmid replication origins

DATE-ISSUED: March 27, 2001

US-CL-CURRENT: 435/6; 435/252.3, 435/320.1, 435/471, 435/91.1, 536/23.1, 536/24.1

APPL-NO: 9/ 134246 DATE FILED: August 14, 1998 IN: Wayne; Jay, Xu; Shuang-yong

AB: The present invention relates to cloned DNA containing origin of DNA replication and to cloned DNA encoding repliation protein, RepT.

L6: Entry 4 of 7

File: USPT

Mar 27, 2001

DOCUMENT-IDENTIFIER: US 6207377 B1 TITLE: Method for construction of thermus-E. coli shuttle vectors and identification of two Thermus plasmid replication origins

Brief Summary Paragraph Right (1):
The present invention relates to recombinant DNA molecules encoding plasmid DNA replication origins in Thermus, as well as to shuttle vectors which contain the same.

Brief Summary Paragraph Right (4):
A Thermus-E. coli shuttle vector would be desirable if one needs to have the convenience of cloning in E. coli, isolation of DNA from E. coli for further manipulations and subsequently gene selection and expression in Thermus. Such Thermus-E. coli shuttle vectors could be used to screen, select and express thermostable proteins in Thermus. Using these vectors, a gene could, for example, be mutated within a mesophile, transferred to a thermophile, and then its encoded protein selected for increased thermostability. In this way, mesophile-thermophile shuttle-vectors can be used to conduct directed evolution, or protein engineering, on desirable gene products.

Brief Summary Paragraph Right (6):
The present invention relates to recombinant DNA molecules encoding plasmid DNA replication origins in Thermus, as well as to

shuttle vectors which contain the same.

Detailed Description Paragraph Right (17):
The repeats and inverted repeats are important for pTsp45L origin of replication, because deletion of these repeats in a HindIII fragment abolished DNA replication in Thermus. The DNA sequence of pTsp45L is shown in FIG. 7. The Thermus-E. coli shuttle vector containing pTsp45L DNA replication origin was named as pUC-EKR-Tsp45L9Kb.

Detailed Description Paragraph Type 1 (8): 8. To reduce the size of the Thermus replication origin, the 4.2 kb XbaI fragment was further digested with restriction enzymes and subcloned into pUC-EKF or pUC-EKR. One recombinant plasmid contained a 2.3 kb NheI fragment that replicates in Thermus and E. coli. This plasmid pUC-EKF-Tsp3 is a Thermus-E. coli shuttle vector.

Detailed Description Paragraph Type 1 (9):

9. One open reading frame of 1026 bp encoding a 341-amino acid protein was found within the Thermus origin. Deletion of 234 bp (78 amino acid residues) within this gene abolished the Thermus replication function. Insertion of stop codons within this gene causes premature termination and negates the Thermus transformation. Therefore it was determined that this gene (repT) is required for plasmid replication in Thermus HB27 (Pro.sup.-) cells.

Detailed Description Paragraph Type 1 (10):

10. Two Thermus promoters were found upstream of the repT gene that are important for repT expression.

5. Document ID: US 5872238 A

L6: Entry 5 of 7

File: USPT

Feb 16, 1999

US-PAT-NO: 5872238

DOCUMENT-IDENTIFIER: US 5872238 A

TITLE: Thermophile gene transfer

DATE-ISSUED: February 16, 1999

US-CL-CURRENT: 536/23.7

APPL-NO: 8/912794

DATE FILED: August 18, 1997

PARENT-CASE:

This application is a continuation of application Ser. No. 08/496,932, filed on Jun. 30, 1995, now abandoned which is a continuation-in-part of U.S. patent application Ser. No. 08/265,522 filed Jun. 24, 1994, now abandoned.

IN: Weber; J. Mark, Demirjian; David C., Casadaban; Malcolm J., Vonstein; Veronika, Pagratis; Nikos C.

AB: We have developed a new gene transfer system for extreme thermophiles of the genus Thermus, including

Thermus flavus., using a chromosomal gene, and a thermostable derivative of the kanamycin-resistance gene

(kan.sup.tr2). A plasmid mediated gene-replacement process is used to insert it into the chromosome resulting in the

production of Leu.sup.- Km.sup.r transformants. This system not only allows stable, single-copy gene insertion into

the chromosome of an extreme thermophile, but can be used in the thermo-genetic process described here to generate

thermo-stabilized enzymes and proteins for industrial processes. This host-vector environment makes it possible to

generate further thermo-stabilizing mutations in the kan gene beyond those levels previously reported.

L6: Entry 5 of 7

File: USPT

Feb 16, 1999

DOCUMENT-IDENTIFIER: US 5872238 A

TITLE: Thermophile gene transfer

Brief Summary Paragraph Right (9):

Koyama et al. (1990) A plasmid vector for an extreme thermophile, Thermus thermophilus, FEMS Microbiology Letters 72:97-102, teach a Thermus-E. coli shuttle vector carrying a tryptophan synthetase gene (trpB). This cryptic plasmid pTTS, was able to transform Thermus thermophilus. The authors point out that a plasmid vector carrying trpBA was not suitable for selection since the cloned DNA fragment recombined with the chromosomal counterpart at high frequency.

Brief Summary Paragraph Right (17):

Lasa et al. (1992a) Development of Thermus-Escherichia Shuttle Vectors and Their Use for Expression of the Clostridium thermocellum celA Gene in Thermus thermophilus, J. of Bacteriology 174:6424-6431, teach the self-selection of undescribed origins of replication from cryptic plasmids from uncharacterized Thermus spp. and Thermus aquaticus are isolated and cloned into E. coli vectors. Plasmids were constructed with these origins, pLU1 to pLU4 from T. aquaticus, and pMY1 to pMY3 from Thermus spp. The plasmids then had a modified form of the cellulase gene (celA) from Clostridium thermocellum and were expressed in E. coli with the signal peptide from the S-layer gene from T. thermophilus. Transformation back into T. thermophilus allowed for expression at 70.degree. C.

Other Reference Publication (11):

Lasa et al. (1992a) Development of Thermus-Escherichia Shuttle Vectors and Their Use for Expression of the Clostridium thermocellum celA Gene in Thermus thermophilus, J. of Bacteriology 174:6424-6431.

6. Document ID: US 5786174 A

L6: Entry 6 of 7

File: USPT

Jul

28, 1998

US-PAT-NO: 5786174

DOCUMENT-IDENTIFIER: US 5786174 A

TITLE: Thermophile gene transfer

DATE-ISSUED: July 28, 1998

US-CL-CURRENT: 435/69.1; 435/463, 530/350, 536/23.1

APPL-NO: 8/790309 DATE FILED: January 28, 1997

PARENT-CASE:

This application is a continuation of application Ser. No. 08/265,522, filed on 24 Jun., 1994, now abandoned.

IN: Weber; J. Mark, Demirjian; David C., Casadaban; Malcolm J., Pagratis; Nikos C., Vonstein; Veronika

AB: We have developed a new gene transfer system for extreme thermophiles of the genus Thermus, including

Thermus flavus, using a chromosomal gene, and a thermostable derivative of the kanamycin-resistance gene

(kan.sup.tr2). A plasmid mediated gene-replacement process is used to insert it into the chromosome resulting in the

production of Leu.sup.- Km.sup.r transformants. This system not only allows stable, single-copy gene insertion into

the chromosome of an extreme thermophile, but can be used in the

thermo-genetic process described here to generate thermo-stabilized enzymes and proteins for industrial processes. This host-vector environment makes it possible to

generate further thermo-stabilizing mutations in the kan gene beyond those levels previously reported.

L6: Entry 6 of 7

File: USPT

Jul

28, 1998

DOCUMENT-IDENTIFIER: US 5786174 A TITLE: Thermophile gene transfer

Brief Summary Paragraph Right (10):

Koyama et al. (1990) A plasmid vector for an extreme thermophile, Thermus thermophilus, FEMS Microbiology Letters 72:97-102, teach a Thermus-E. coli shuttle vector carrying a tryptophan synthetase gene (trpB). This cryptic plasmid pTT8, was able to transform Thermus thermophilus. The authors point out that a plasmid vector carrying trpBA was not suitable for selection since the cloned DNA fragment recombined with the chromosomal counterpart at high frequency.

Brief Summary Paragraph Right (18):

Lasa et al. (1992a) Development of Thermus-Escherichia Shuttle Vectors and Their Use for Expression of the Clostridium thermocellum celA Gene in Thermus thermophilus, J. of Bacteriology 174:6424-6431, teach the self-selection of undescribed origins of replication from cryptic plasmids from uncharacterized Thermus spp. and Thermus aquaticus are isolated and cloned into E. coli vectors. Plasmids were constructed with these origins, pLU1 to pLU4 from T. aquaticus, and pMY1 to pMY3 from Thermus spp. The plasmids then had a modified form of the cellulase gene (celA) from Clostridium thermocellum and were expressed in E. coli with the signal peptide from the S-layer gene from T. thermophilus. Transformation back into T. thermophilus allowed for expression at 70.degree. C.

Other Reference Publication (18):

Lasa et al. (1992a) Development of Thermus-Escherichia Shuttle Vectors and Their Use for Expression of the Clostridium thermocellum celA Gene in Thermus thermophilus, J. of Bacteriology 174:6424-6431.

7. Document ID: US 5120658 A

L6: Entry 7 of 7

File: USPT

Jun 9, 1992

US-PAT-NO: 5120658 DOCUMENT-IDENTIFIER: US 5120658 A

TITLE: Thermostable tryptophan synthetase gene and extremely thermophilic plasmid vector incorporating said gene

DATE-ISSUED: June 9, 1992

US-CL-CURRENT: 435/320.1; 435/108, 435/183, 435/252.3, 435/69.1, 435/71.2, 435/91.41, 536/23.2

APPL-NO: 7/ 329765 DATE FILED: March 28, 1989 FOREIGN-APPL-PRIORITY-DATA: COUNTRY

APPL-NO

APPL-DATE

JP

63-163779

June 30, 1988

IN: Koyama; Yoshinori, Furukawa; Kensuke, Tomizuka; Noboru

AB: A DNA segment, specifically a thermostable tryptophan synthetase gene originating in the strain of

extremely thermophilic Thermus aquaticus T2, characterized by the restriction enzyme map of FIG. 1, and not cleaved

by specific restriction enzymes., An extremely thermophilic plasmid vector pYK 105, having the DNA segment and an

Escherichia coli plasmid vector pUC 13 incorporated in a cryptic plasmid pTT8.

L6: Entry 7 of 7

File: USPT

Jun 9, 1992

DOCUMENT-IDENTIFIER: US 5120658 A

TITLE: Thermostable tryptophan synthetase gene and extremely thermophilic plasmid vector incorporating said gene

Detailed Description Paragraph Right (23):

The plasmid pYK 105 separated from the transformed strain possesses the structure illustrated at the bottom of FIG. 4. Since it possesses the pUC 13 plasmid, it constitutes a shuttle vector which can replicate not only in the microorganism of genus Thermus but also in the Escherichia coli. The selection of the transformed strain is attained by virtue of the tolerance to Ampicillin in the case of the Escherichia coli and the complementation of the tryptophan-demanding property in the case of the thermophilus strain of Thermus thermophilus HB 27 trp.sup. The pYK 105 is the first selectable plasmid vector produced with a microorganism of genus Thermus.

Detailed Description Paragraph Right (59):

When this culture was continued at 70.degree. C. for two days, there was obtained a transformed strain of Thermus thermophilus HB 27 trp.sup.- (pyK 105) no longer demanding tryptophan. The plasmid pyK 105 separated from the transformed strain possessed the structure illustrated at the bottom of FIG. 4. Since it possessed a pUC 13 plasmid, it constituted a shuttle vector which can replicate not only in the microorganism of genus Thermus but also in the Escherichia coli. The selection of the transformed strain could be effected by virtue of the ampicillin resistance in the case of the Escherichia coli and by virtue of the complementation of the tryptophan-demanding property in the case of the thermophilus HB 27 trp.sup.-. The pYK 105 is the first selectable plasmid vector constructed for the microorganism of genus Thermus.

09/664186 ANT PG

=> s thermus

8444 THERMUS L1

=> s shuttle

33438 SHUTTLE L2

=> s rept

1317 REPT L3

=> s promoter

L4 385651 PROMOTER

=> s 11 and 12

46 L1 AND L2

=> dup rem 15

PROCESSING COMPLETED FOR L5

18 DUP REM L5 (28 DUPLICATES REMOVED)

=> d 16 ibib abs 1-18

L6 ANSWER I OF 18 WPIDS (C) 2002 THOMSON DERWENT

ACCESSION NUMBER: 2002-269188 [31] WPIDS

DOC. NO. CPI:

C2002-079914

TITLE:

Producing transformed microorganism, preferably Bacillus, involves selecting competent microorganism, producing DNA

construct in vitro, and directly transforming the microorganism with the DNA construct.

DERWENT CLASS: B04 D16

DIAZ-TORRES, M R; LEE, E W; MORRISON, T B; INVENTOR(S):

SCHELLENBERGER, V; SELIFONOVA, O V

PATENT ASSIGNEE(S): (GEMV) GENENCOR INT INC

COUNTRY COUNT: PATENT INFORMATION:

PATENT NO KIND DATE WEEK LA PG

WO 2002014490 A2 20020221 (200231)* EN 48

RW: AT BE CH CY DE DK EA ES FI FR GB GH GM GR IE IT KE LS EU MC MW MZ

NL OA PT SD SE SL SZ TR TZ UG ZW

W: AE AG AL AM AT AU AZ BA BB BG BR BY BZ CA CH CN CO CR CU CZ DE DK

DM DZ EC EE ES FI GB GD GE GH GM HR HU ID IL IN IS JP KE KG KP KR

KZ LC LK LR LS LT LU LV MA MD MG MK MN MW MX MZ NO NZ PL PT RO RU

SD SE SG SI SK SL TJ TM TR TT TZ UA UG UZ VN YU ZA ZW

APPLICATION DETAILS:

PATENT NO KIND

APPLICATION DATE

WO 2002014490 A2

WO 2001-US25166 20010810

PRIORITY APPLN. INFO: US 2000-224948P 20000811 AN 2002-269188 [31] WPIDS

AB WO 200214490 A UPAB: 20020516

NOVELTY - Producing (M1) a transformed microorganism, preferably

involves selecting a competent microorganism, producing a DNA construct in

vitro, and directly transforming the microorganism with the DNA construct

such that the DNA construct becomes integrated into a chromosome of the microorganism.

DETAILED DESCRIPTION - INDEPENDENT CLAIMS are also included for the

(1) a library of mutants (I) produced by M1;

(2) directed evolution (M2) of a sequence in the host cell chromosome, involves in vitro mutagenesis of a DNA construct, direct transformation of the mutagenized sequence into a competent host cell, screening for, or selection of, mutants possessing or exhibiting a desired property, and repeating the above mentioned steps for one or more rounds;

(3) constructing (M3) a sequence of interest at a target sequence of a selected microorganism, where the target sequence includes a residing marker, involves assembling a DNA construct in vitro comprising an incoming sequence, a selectable marker, and two flanking sequences

are homologous to sequences of the target sequence, where the selectable marker of the DNA construct is different than the residing marker of the microorganism, transforming the microorganism with the DNA construct

conditions permitting the incoming sequence and selectable marker to inactivate the residing marker, and selecting for transformants that include the selectable marker, and repeating the above mentioned steps, where with each repetition of the DNA construct comprises a selectable marker different from the selectable marker in the previous step and the selectable marker of the previous step acts as the residing marker in the microorganisms.

USE - M1 is useful for producing a transformed microorganisms selected from Acinetobacter, ***Thermus*** , Deinococcus, Radiodurans.

and Bacillus, preferably Bacillus, where the Bacillus is a super-competent strain, preferably a Pxyl-comK strain (claimed).

ADVANTAGE - M1 transforms the DNA constructs into the

with good efficiency, and allows for the generation of large libraries. M1 provides in vitro construction and direct transformation into Bacillus, without the use of any intervening microorganisms. M1 does not require antibiotic or other selectable marker to maintain the plasmid in the cells, which is undesirable for production strains and constrains choice of screening conditions. M1 allows evolution of single copy genes of a strain, and prevents variations in copy number which skews a library. Dwg.0/15

L6 ANSWER 2 OF 18 MEDLINE

ACCESSION NUMBER: 2002255802 IN-PROCESS DOCUMENT NUMBER: 21911539 PubMed ID: 11914489

Comparative analysis of space-grown and earth-grown TITLE: crystals of an aminoacyl-tRNA synthetase: space-grown

crystals are more useful for structural determination.

AUTHOR: Ng Joseph D; Sauter Claude; Lorber Bernard; Kirkland Natalie; Arnez John; Giege Richard

CORPORATE SOURCE: Departement Mecanismes et Macromolecules de la Synthese

Proteique et Cristallogenese, UPR 9002, Institut de Biologie Moleculaire et Cellulaire du CNRS, 15 Rue Rene

Descartes, F-67084 Strasbourg CEDEX, France. ACTA CRYSTALLOGRAPHICA. SECTION D: SOURCE:

BIOLOGICAL

CRYSTALLOGRAPHY, (2002 Apr) 58 (Pt 4) 645-52.

Journal code: 9305878. ISSN: 0907-4449.

PUB. COUNTRY: Denmark

Journal; Article; (JOURNAL ARTICLE) English

LANGUAGE:

IN-PROCESS; NONINDEXED; Priority Journals FILE SEGMENT: PDB-110W; PDB-R110WSF

OTHER SOURCE: ENTRY DATE:

Entered STN: 20020509

Last Updated on STN: 20020509

AB Protein crystallization under microgravity aims at benefiting from the quasi-absence of convection and sedimentation to favor well ordered crystal nucleation and growth. The dimeric multidomain enzyme aspartyl-tRNA synthetase from ***Thermus*** thermophilus has been crystallized within dialysis reactors of the Advanced Protein Crystallization Facility in the laboratory on earth and under microgravity aboard the US Space ***Shuttle*** . A strictly comparative crystallographic analysis reveals that the crystals grown in space are superior in every respect to control crystals prepared in otherwise identical conditions on earth. They diffract X-rays more intensely and have a lower mosaicity, facilitating the process of protein structure determination. Indeed, the electron-density map calculated from diffraction data of space-grown crystals contains considerably more detail. The resulting three-dimensional structure model at 2.0 A resolution is more accurate than that produced in parallel using the data originating from earth-grown crystals. The major differences between the structures, including the better defined amino-acid side chains and the higher order of bound water molecules, are emphasized.

L6 ANSWER 3 OF 18 HCAPLUS COPYRIGHT 2002 ACS ACCESSION NUMBER: 2001:221921 HCAPLUS

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Replication origins and proteins of plasmids of the thermophilic bacterium ***Thermus*** and the
                                                                                      ABSTRACTS INC.DUPLICATE
TITLE:
                                                                                        3
                                                                                      ACCESSION NUMBER: 2001:186597 BIOSIS
              construction of ***Thermus*** -E. coli
                                                                                      DOCUMENT NUMBER: PREV200100186597
               ***shuttle*** vectors
                                                                                                     Characterization of the minimal replicon of a cryptic
                                                                                      TITLE:
INVENTOR(S): Wayne, Jay; Xu, Shuang-Yong
PATENT ASSIGNEE(S): New England Biolabs, Inc., USA
SOURCE: U.S., 32 pp.
                                                                                                 Deinococcus radiodurans SARK plasmid and development of
                                                                                                  versatile Escherichia coli-D. radiodurans ***shuttle***
                                                                                                  vectors.
              CODEN: USXXAM
                                                                                                         Meima, Rob; Lidstrom, Mary E. (1)
                                                                                      AUTHOR(S):
DOCUMENT TYPE:
                          Patent
                                                                                      CORPORATE SOURCE: (1) Department of Chemical Engineering,
LANGUAGE:
                       English
                                                                                      University of
FAMILY ACC. NUM. COUNT: 1
                                                                                                  Washington, Seattle, WA, 98195-1750:
PATENT INFORMATION:
                                                                                                  lidstrom@u.washington.edu USA
                                                                                                        Applied and Environmental Microbiology, (September,
                                                                                      SOURCE:
   PATENT NO. KIND DATE
                                        APPLICATION NO. DATE
                                                                                      2000)
   US 6207377 B1 20010327 US 1998-134246 19980814
                                                                                                  Vol. 66, No. 9, pp. 3856-3867. print.
                                                                                                  ISSN: 0099-2240.
AB Replication origins and functions of two plasmids of the thermophilic
                                                                                      DOCUMENT TYPE: Article
   bacterium ***Thermus*** YS45 are described. Two genes, oriT of
                                                                                      LANGUAGE:
                                                                                                          English
   pTsp45S and parA of pTsp45L, that are essential for replication are
                                                                                      SUMMARY LANGUAGE: English
cloned
                                                                                      AB The nucleotide sequence of a 12-kb fragment of the cryptic Deinococcus
   and characterized. These functions may be useful in the construction of
                                                                                         radiodurans SARK plasmid pUE10 was determined, in order to direct the
    ***Thermus*** -Escherichia coli ***shuttle*** vectors.
                                                                                         development of small, versatile cloning systems for Deinococcus.
                            20 THERE ARE 20 CITED REFERENCES
REFERENCE COUNT:
                                                                                         Annotation of the sequence revealed 12 possible open reading frames.
AVAILABLE FOR THIS
                  RECORD. ALL CITATIONS AVAILABLE IN THE RE
                                                                                         these are the repU and resU genes, the predicted products of which share
FORMAT
                                                                                         similarity with replication proteins and site-specific resolvases,
                                                                                         respectively. The products of both genes were demonstrated using an
L6 ANSWER 4 OF 18 BIOSIS COPYRIGHT 2002 BIOLOGICAL
                                                                                         overexpression system in Escherichia coli. RepU was found to be required
ABSTRACTS INC.DUPLICATE
                                                                                         for replication, and ResU was found to be required for stable maintenance
                                                                                         of pUE10 derivatives. Gel shift analysis using purified His-tagged RepU
ACCESSION NUMBER: 2001:462444 BIOSIS DOCUMENT NUMBER: PREV200100462444
                                                                                         identified putative binding sites and suggested that RepU may be involved
                                                                                         in both replication initiation and autoregulation of repU expression. In
               Production of recombinant alpha-galactosidases in
TITLE:
                                                                                         addition, a gene encoding a possible antirestriction protein was found,
              ***Thermus*** thermophilus.
                                                                                         which was shown to be required for high transformation frequencies. The
                   Fridjonsson, Olafur (1); Mattes, Ralf
 AUTHOR(S):
                                                                                         arrangement of the replication region and putative replication genes for
 CORPORATE SOURCE: (1) Prokaria Ltd., Gylfaflot 5, 112, Reykjavik:
                                                                                         this plasmid from D. radiodurans strain SARK is similar to that for plasmids found in ***Thermus*** but not to that for the 45.7-kb
            olafur@prokaria.com Iceland
                  Applied and Environmental Microbiology, (September,
 SOURCE:
                                                                                         plasmid found in D. radiodurans strain R1. The minimal region required
 2001)
                                                                                       for
             Vol. 67, No. 9, pp. 4192-4198. print.
                                                                                         autonomous replication in D. radiodurans was determined by sequential
            ISSN: 0099-2240.
                                                                                         deletion of segments from the 12-kb fragment. The resulting minimal
 DOCUMENT TYPE: Article
                                                                                         replicon, which consists of approximately 2.6 kb, was used for the
                    English
 LANGUAGE:
                                                                                          construction of a ***shuttle*** vector for E. coli and D. radiodurans.
 SUMMARY LANGUAGE: English
 AB A ***Thermus*** thermophilus selector strain for production of
                                                                                          This vector, pRAD1, is a convenient general-purpose cloning vector. In
                                                                                          addition, pRAD1 was used to generate a promoter probe vector, and a
   thermostable and thermoactive alpha-galactosidase was constructed. For
                                                                                         plasmid containing lacZ and a Deinococcus promoter was shown to
    this purpose, the native alpha-galactosidase gene (agaT) of T.
                                                                                         efficiently express LacZ.
    thermophilus TH125 was inactivated to prevent background activity. In
 our
                                                                                       L6 ANSWER 6 OF 18 BIOSIS COPYRIGHT 2002 BIOLOGICAL
   first attempt, insertional mutagenesis of agaT by using a cassette
                                                                                       ABSTRACTS INC.DUPLICATE
   carrying a kanamycin resistance gene led to bacterial inability to utilize
    melibiose (alpha-galactoside) and galactose as sole carbohydrate sources
                                                                                       ACCESSION NUMBER: 1999:199329 BIOSIS
    due to a polar effect of the insertional inactivation. A Gal+ phenotype
                                                                                       DOCUMENT NUMBER: PREV199900199329
    was assumed to be essential for growth on melibiose. In a Gal-
                                                                                                  Dissimilatory reduction of Fe(III) and other electron acceptors by a ***Thermus*** isolate.
                                                                                       TITLE:
 background.
    accumulation of galactose or its metabolite derivatives produced from
                                                                                       AUTHOR(S):
                                                                                                          Kieft, T. L. (1); Fredrickson, J. K.; Onstott, T. C.;
    melibiose hydrolysis could interfere with the growth of the host strain
                                                                                                   Gorby, Y. A.; Kostandarithes, H. M.; Bailey, T. J.;
    harboring recombinant alpha-galactosidase. Moreover, the AgaT- strain
                                                                                                   Kennedy, D. W.; Li, S. W.; Plymale, A. E.; Spadoni, C. M.;
 had
                                                                                                   Gray, M. S.
    to be Kms for establishment of the plasmids containing
                                                                                       CORPORATE SOURCE: (1) Department of Biology, New Mexico Institute
 alpha-galactosidase
                                                                                       of Mining
    genes and the kanamycin resistance marker. Therefore, a suitable selector
                                                                                                   and Technology, Socorro, NM, 87801 USA
    strain (AgaT- Gal+ Kms) was generated by applying integration
                                                                                       SOURCE:
                                                                                                         Applied and Environmental Microbiology, (March, 1999)
 mutagenesis
                                                                                       Vol.
    in combination with phenotypic selection. To produce heterologous
                                                                                                   65, No. 3, pp. 1214-1221.
    alpha-galactosidase in T. thermophilus, the isogenes agaA and agaB of
                                                                                                   ISSN: 0099-2240.
    Bacillus stearothermophilus KVE36 were cloned into an Escherichia coli-
     ***Thermus*** ***shuttle*** vector. The region containing the E.
                                                                                       DOCUMENT TYPE: Article
    coli plasmid sequence (pUC-derived vector) was deleted before
                                                                                                           English
                                                                                       LANGUAGE:
                                                                                       AB A thermophilic bacterium that can use O2, NO3-, Fe(III), and S0 as
    transformation of T. thermophilus with the recombinant plasmids. As a
                                                                                          terminal electron acceptors for growth was isolated from groundwater
    result, transformation efficiency and plasmid stability were improved.
                                                                                          sampled at a 3.2-km depth in a South African gold mine. This organism,
    However, growth on minimal agar medium containing melibiose was
                                                                                          designated SA-01, clustered most closely with members of the genus
                                                                                           ***Thermus***, as determined by 16S rRNA gene (rDNA) sequence
    only following random selection of the clones carrying a plasmid-based
    mutation that had promoted a higher copy number and greater stability of
                                                                                          The 16S rDNA sequence of SA-01 was >98% similar to that of
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134:247953

DOCUMENT NUMBER:

the plasmid.

1.6 ANSWER 5 OF 18 BIOSIS COPYRIGHT 2002 BIOLOGICAL

strain NMX2 A.1, which was previously isolated by other investigators from

a thermal spring in New Mexico. Strain NMX2 A.1 was also able to

Fe(HI) and other electron acceptors. Neither SA-01 nor NMX2 A.1 grew fermentatively, i.e., addition of an external electron acceptor was required for anaerobic growth. ***Thermus*** strain SA-01 reduced soluble Fe(III) complexed with citrate or nitrilotriacetic acid (NTA); however, it could reduce only relatively small quantities (0.5 mM) of hydrous ferric oxide except when the humic acid analog 2,6-anthraquinone disulfonate was added as an electron ***shuttle***, in which case 10 mM Fe(III) was reduced. Fe(III)-NTA was reduced quantitatively to

reduction of Fe(III)-NTA was coupled to the oxidation of lactate and supported growth through three consecutive transfers. Suspensions of ***Thermus*** strain SA-01 cells also reduced Mn(IV),

Co(III)-EDTA. Cr(VI), and U(VI). Mn(IV)-oxide was reduced in the presence of either lactate or H2. Both strains were also able to mineralize NTA to CO2 and

couple its oxidation to Fe(III) reduction and growth. The optimum temperature for growth and Fe(III) reduction by ***Thermus*** strains SA-01 and NMX2 A.1 is approximately 65degreeC; their optimum pH is 6.5 to

7.0. This is the first report of a ***Thermus*** sp. being able to couple the oxidation of organic compounds to the reduction of Fe, Mn, or

L6 ANSWER 7 OF 18 BIOSIS COPYRIGHT 2002 BIOLOGICAL ABSTRACTS INC.DUPLICATE

ACCESSION NUMBER: 2000:51935 BIOSIS

DOCUMENT NUMBER: PREV200000051935

Ribosomal gene disruption in the extreme thermophile TITLE: ***Thermus*** thermophilus HB8. Generation of a mutant

lacking ribosomal protein \$17. AUTHOR(S): Simitsopoulou, Maria; Avila, Horacio; Franceschi,

Francois

CORPORATE SOURCE: (1) Max-Planck-Institut fuer Molekulare Genetik,

Ribosomen, Ihnestrasse 73, Berlin, 14195 Germany European Journal of Biochemistry, (Dec., 1999) Vol. 266, SOURCE: No. 2, pp. 524-532.

ISSN: 0014-2956. DOCUMENT TYPE: Article

LANGUAGE: English SUMMARY LANGUAGE: English

AB S17 is a primary rRNA-binding protein which has been implicated in ribosome assembly and translational fidelity. We describe the generation and biochemical characterization of an S17 minus ribosomal mutant, a ribosomal protein-lacking mutant obtained in ***Thermus** thermophilus HB8. The S17 mutant was obtained by insertional inactivation

of the target gene with the kanamycin adenyl transferase (kat) gene, making use of a ***Thermus*** -Escherichia ***shuttle*** vector

the natural ability of ***Thermus*** to transform. In the final construct used to transform ***Thermus*** cells, the S17 coding region was replaced with the kat gene cloned in-frame with the first three amino acids of S17. Hence, in vivo transcription of the kat gene was under the control of the ribosomal operon promoter. As in Escherichia coli, the ***Thermus*** S17 mutant exhibited a temperature-sensitive

Two-dimensional PAGE, Western blot, and ELISA confirmed the absence

from the mutant ribosomes. Sucrose-gradient profiles of mutant cells showed a clear separation and normal proportions of 50S and 30S subunits and a normal ratio between them. In addition, the S17 mutant showed the presence of a 20S peak representing assembly-defective particles. The successful re-incorporation of protein S17 into the mutant ribosomes was demonstrated when reconstitution with isolated S17 was performed at 60 degreeC.

L6 ANSWER 8 OF 18 BIOSIS COPYRIGHT 2002 BIOLOGICAL ABSTRACTS INC.DUPLICATE

ACCESSION NUMBER: 2000:27289 BIOSIS

DOCUMENT NUMBER: PREV200000027289

A high-transformation-efficiency cloning vector for TITLE: ***Thermus*** thermophilus.

de Grado, Myriam; Castan, Pablo; Berenguer, Jose (1) AUTHOR(S): CORPORATE SOURCE: (1) Centro de Biologa Molecular "Severo Ochoa", UAM-CSIC,

Universidad Autonoma de Madrid, 28049, Madrid Spain SOURCE: Plasmid, (Nov., 1999) Vol. 42, No. 3, pp. 241-245.

ISSN: 0147-619X. DOCUMENT TYPE: Article LANGUAGE: English SUMMARY LANGUAGE: English

AB The cloning vector pMK18 was developed through the fusion of the minimal

replicative region from an indigenous plasmid of ***Thermus*** sp. ATCC27737, a gene cassette encoding a thermostable resistance to kanamycin, and the replicative origin and multiple cloning site of pUC18. Plasmid pMK18 showed transformation efficiencies from 108 to 109 per microgram of plasmid in ***Thermus*** thermophilus HB8 and HB27, both

by natural competence and by electroporation. We also show that T. thermophilus HB27 can take pMK18 modified by the Escherichia coli methylation system with the same efficiency as its own DNA. To

its usefulness as a cloning vector, a gene encoding the beta-subunit of a thermostable nitrate reductase was directly cloned in T. thermophilus HB27

from a gene library. Its further transfer to E. coli also proved its utility as a ***shuttle*** vector.

L6 ANSWER 9 OF 18 BIOSIS COPYRIGHT 2002 BIOLOGICAL ABSTRACTS INC.DUPLICATE

ACCESSION NUMBER: 1997:414348 BIOSIS DOCUMENT NUMBER: PREV199799706391

A new ***Thermus*** -Escherichia coli ***shuttle*** TITLE: integration vector system.

Tamakoshi, Masatada; Uchida, Manabu; Tanabe, AUTHOR(S):

Kazuhiro:

Fukuyama, Shiro; Yamagishi, Akihiko (1); Oshima, Tairo CORPORATE SOURCE: (1) Dep. Mol. Biol., Tokyo Univ. Pharmacy Life Sci., 1432

Horinouchi, Hachioji, Tokyo 192-03 Japan Journal of Bacteriology, (1997) Vol. 179, No. 15, pp.

SOURCE: 4811-4814

ISSN: 0021-9193

DOCUMENT TYPE: Article English LANGUAGE:

AB We established a ***Thermus*** thermophilus strain in which the ругЕ

gene (coding for orotate phosphoribosyltransferase of the pyrimidine biosynthetic pathway) was totally deleted. We also constructed an integration vector, which consisted of the Escherichia coli plasmid vector pBluescript and a 2.1-kb segment of the T. thermophilus leu operon sequence, for the integration of a foreign gene into a chromosome of the thermophile, pyrE and leuB genes were used as probes to test the integration vector. The integration vector pINV, bearing the pyrE gene, transformed the DELTA-pyrE strain at a frequency of 6 times 10-5 through a

single crossover event. The leuB gene could also be used as another marker

of the integration vector system. The vector could be integrated at the expected site. By digesting the chromosomal DNA of the T. thermophilus transformants with a unique restriction enzyme, the vector could be recovered into E. coli after the recircularization in vitro. The kanamycin nucleotidyltransferase gene could be successfully expressed in the thermophile by using pINV.

L6 ANSWER 10 OF 18 BIOSIS COPYRIGHT 2002 BIOLOGICAL ABSTRACTS INC.DUPLICATE

ACCESSION NUMBER: 1997:506817 BIOSIS DOCUMENT NUMBER: PREV199799806020

Lumenal proteins involved in respiratory electron transport TITLE: in the cyanobacterium Synechocystis sp. PCC6803.

AUTHOR(S): Manna, Pradip; Vermaas, Wim (1)

CORPORATE SOURCE: (1) Molecular Cellular Biol. Program, Dep. Botany, Cent.

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Study Early Events Photosynthesis, Arizona State Univ., Box
            871601, Tempe, AZ 85287-1601 USA
                  Plant Molecular Biology, (1997) Vol. 35, No. 4, pp.
SOURCE:
            407-416.
            ISSN: 0167-4412.
DOCUMENT TYPE:
                       Article
LANGUAGE:
                     English
AB Cyanobacterial thylakoids catalyze both photosynthetic and respiratory
   activities. In a photosystem I-less Synechocystis sp. PCC 6803 strain,
   electrons generated by photosystem II appear to be utilized by cytochrome
   oxidase. To identify the lumenal electron carriers (plastocyanin and/or
   cytochromes c-553, c-550, and possibly c-M) that are involved in transfer
   of photosystem II-generated electrons to the terminal oxidase, deletion
   constructs for genes coding for these components were introduced into a
   photosystem I-less Synechocystis sp. PCC 6803 strain, and electron flow
   out of photosystem II was monitored in resulting strains through
   chlorophyll fluorescence yields. Loss of cytochrome c-553 or
plastocyanin,
   but not of cytochrome c-550, decreased the rate of electron flow out of
   photosystem II. Surprisingly, cytochrome c-M could not be deleted in a
   photosystem I-less background strain, and also a double-deletion mutant
   lacking both plastocyanin and cytochrome c-553 could not be obtained.
   Cytochrome c-M has some homology with the cytochrome c-binding
regions of
   the cytochrome caa-3-type cytochrome oxidase from Bacillus spp. and
***Thermus*** thermophilus. We suggest that cytochrome c-M is a
   component of cytochrome oxidase in cyanobacteria that serves as redox
   intermediate between soluble electron carriers and the cytochrome aa-3
   complex, and that either plastocyanin or cytochrome c-553 can
     ***shuttle*** electrons from the cytochrome b-6f complex to
cytochrome
   c-M.
L6 ANSWER 11 OF 18 BIOSIS COPYRIGHT 2002 BIOLOGICAL
ABSTRACTS INC.DUPLICATE
 ACCESSION NUMBER: 1997:453208 BIOSIS
DOCUMENT NUMBER: PREV199799752411
 TITLE:
                 Identification of a thermophilic plasmid origin and its
             cloning within a new ***Thermus*** -E. coli
              ***shuttle*** vector.
                     Wayne, Jay; Xu, Shuang-Yong (1)
 AUTHOR(S):
 CORPORATE SOURCE: (1) New Engl. Biolabs, 32 Tozer Road, Beverly,
 MA 01915 USA
                   Gene (Amsterdam), (1997) Vol. 195, No. 2, pp. 321-328.
 SOURCE:
             ISSN: 0378-1119.
 DOCUMENT TYPE: Article
 LANGUAGE:
                      English
 AB A pUC19-based vector has been generated for selecting functional
    thermophilic origins (oris) of ***Thermus*** ssp. Once combined with
    thermophilic DNA, the vector can be amplified in ampicillin resistant
    (Ap-R) E. coli, prior to transformation and kanamycin (Km) selection in
     ***Thermus*** thermophilus. The Km-R ***Thermus***
 transformants
    replicate any newly-formed ***shuttle*** vectors via introduced thermophilic oris. Using this "ori-selecting" vector, three novel
    thermophilic oris were cloned from randomly digested ***Thermus***
cryptic plasmid DNA. These ***shuttle*** vectors are useful for
    genetic analyses, as well as protein engineering within thermophiles. The
    smallest ori-containing sequence of 4.2 kb has been subcloned, sequenced,
    and further refined to 2.3 kb. A significant ORF of 341 amino acids (aa),
    with a ***Thermus*** promoter and RBS, is found within the
    thermophilic ori. Deleting part of this ORF abolishes the ***shuttle***
    vector's ability to replicate in T. thermophilus. Therefore, we postulate
    that this ORF encodes a replication protein (Rep) necessary for
    thermophilic plasmid replication. The thermophilic ori also contains two
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sequences which resemble DnaA boxes.

L6 ANSWER 12 OF 18 BIOSIS COPYRIGHT 2002 BIOLOGICAL ABSTRACTS INC.DUPLICATE
10
ACCESSION NUMBER: 1993:4035 BIOSIS
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AUTHOR(S): Lasa, Inigo; De Grado, M.; De Pedro, M. A.; Berenguer,

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Jose
CORPORATE SOURCE: (1) Centro de Biologie Molecular, Universidad
Autonoma de
             Madrid-Consejo Superior de Investigaciones Científicas,
             28049 Madrid Spain
                   Journal of Bacteriology, (1992) Vol. 174, No. 20, pp.
SOURCE:
             6424-6431.
            ISSN: 0021-9193.
DOCUMENT TYPE: Article
                     English
LANGUAGE:
AB We describe the self-selection of replication origins of undescribed
   cryptic plasmids from ***Thermus*** aquaticus Y-VII-51B (ATCC
   and a ***Thermus*** sp. strain (ATCC 27737) by random insertion of
   thermostable kanamycin adenyltransferase cartridge. Once selected, these
   autonomous replication origins were cloned into the Escherichia coli
   vector pUC9 or pUC19. The bifunctional plasmids were analyzed for their sizes, relationships, and properties as ***shuttle*** vectors for
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Thermus -Escherichia cloning. Seven different vectors with diverse kanamycin resistance levels, stabilities, transformation efficiencies, and copy numbers were obtained. As a general rule, those from T. aquaticus

(pLU1 to pLU4) were more stable than those from the ***Thermus*** sp.
(pMY1 to pMY3). To probe their usefulness, we used one of the plasmids (pMY1) to clone in E. coli a modified form of the cellulase gene (celA) from Clostridium thermocellum in which the native signal peptide was replaced in vitro by that from the S-layer gene of T. thermophilus HB8. The hybrid product was expressed and exported by E. coli. When the gene was transferred by transformation into T. thermophilus, the cellulase

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protein was also expressed and secreted at 70 degree C.
L6 ANSWER 13 OF 18 MEDLINE
ACCESSION NUMBER: 90363893 MEDLINE
DOCUMENT NUMBER: 90363893 PubMed ID: 2203048
             Molecular structures and evolution of mouse isozyme genes
TITLE:
          functioning in the malate-aspartate ***shuttle***
               Shimada K; Joh T; Ding S H; Choudhury B K; Setoyama
AUTHOR:
CORPORATE SOURCE: Department of Biochemistry, Kumamoto
University Medical
          School, Japan.
SOURCE:
               PROGRESS IN CLINICAL AND BIOLOGICAL
RESEARCH, (1990) 344
          139-58.
          Journal code: PZ5; 7605701. ISSN: 0361-7742.
PUB. COUNTRY:
                   United States
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Journal; Article; (JOURNAL ARTICLE)
LANGUAGE: English
FILE SEGMENT: Priority Journals
ENTRY MONTH: 199010
ENTRY DATE: Entered STN: 19901109
Last Updated on STN: 19980206

Entered Medline: 19901003
AB To examine molecular mechanisms of transcription of mammalian isozyme

genes functioning in the malate-aspartate ***shuttle*** and to observe structural and evolutionary relationships, we investigated gene organizations of cAspAT and mAspAT, and cMDH and mMDH, and sletted and

characterized cDNAs and genomic DNAs for these isozymes in mice. The deduced amino acid sequences of mouse cAspAT and mAspAT showed about 47%,

and those of mouse cMDH and mMDH, about 23% overall homology. Surprisingly, the homology between the mouse cMDH and thermophilic bacterial MDH, as well as the homology between the mouse mMDH and E. coli

MDH, markedly exceeds the intraspecies sequence homology between mMDH and

cMDH from mice. The first duplication of a common ancestral MDH gene should thus have occurred long before the emergence of the eukaryotic cells, and subsequently, the mammalian mMDH and E. coli MDH genes have

evolved from one of the duplicates. The mammalian cMDH and ***Thermus***

flavus MDH genes have no doubt evolved from one of the other

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duplicates.
  Moreover, structural organizations of the two-pairs of isozyme genes
  indicated that introns antedate the divergence of these mitochondrial and
  cytosolic isozyme genes. The 5' ends of all four isozyme genes lacked the
  TATA and CAAT boxes characteristic of eukaryotic promoters but did
  G + C-rich sequences and multiple transcription-initiation sites. We found
   several highly conserved regions in the 5' flanking sequences between
  mAspAT and cAspAT, between mMDH and mAspAT, and between
cMDH and cAspAT
L6 ANSWER 14 OF 18 BIOSIS COPYRIGHT 2002 BIOLOGICAL
ABSTRACTS INC.DUPLICATE
ACCESSION NUMBER: 1991:34771 BIOSIS
DOCUMENT NUMBER: BR40:11751
             A PLASMID VECTOR FOR AN EXTREME
TITLE:
THERMOPHILE ***THERMUS***
          -THERMOPHILUS.
                 KOYAMA Y; ARIKAWA Y; FURUKAWA K
AUTHOR(S):
CORPORATE SOURCE: FERMENTATION RES. INST., AIST, MITI,
TSUKUBA SCI. CITY,
           IBARAKI 305, JAPAN.
                FEMS Microbiol. Lett., (1990) 72 (1-2), 97-102.
           CODEN: FMLED7. ISSN: 0378-1097.
                  BR; OLD
FILE SEGMENT:
LANGUAGE:
                  English
L6 ANSWER 15 OF 18 MEDLINE ACCESSION NUMBER: 91130853 MEDLINE
DOCUMENT NUMBER: 91130853 PubMed ID: 2283046
              A plasmid vector for an extreme thermophile,
             ***Thermus*** thermophilus.
                 Koyama Y; Arikawa Y; Furukawa K
 AUTHOR:
 CORPORATE SOURCE: Fermentation Research Institute, AIST, MITI,
 Tsukuba
           Science City, Ibaraki, Japan.
                FEMS MICROBIOLOGY LETTERS, (1990 Oct) 60 (1-2)
 SOURCE:
 97-101.
           Journal code: FML; 7705721. ISSN: 0378-1097.
                    Netherlands
 PUB. COUNTRY:
           Journal; Article; (JOURNAL ARTICLE)
                   English
 LANGUAGE:
 FILE SEGMENT:
                    Priority Journals
                     199103
 ENTRY MONTH:
                   Entered STN: 19910405
 ENTRY DATE:
           Last Updated on STN: 19910405
           Entered Medline: 19910321
 AB The host-vector system for an extreme thermophile, ***Thermus***
    thermophilus HB27, was developed. The host strain has a mutation in
    tryptophan synthetase gene (trpB), and the mutation was determined to be
    missense mutation by DNA sequence analysis. A ***Thermus*** -E.
 coli
     ***shuttle*** vector pYK109 was constructed. pYK109 consists of
     ***Thermus*** cryptic plasmid pTT8, tryptophan synthetase gene
 (trpB) of
     ***Thermus*** T2 and E. coli plasmid vector pUC13. pYK109
 transformed T.
    thermophilus HB27 trpB5 to Trp+ at a frequency of 10(6) transformants
    microgram DNA.
  L6 ANSWER 16 OF 18 HCAPLUS COPYRIGHT 2002 ACS
                            1990:2020 HCAPLUS
  ACCESSION NUMBER:
  DOCUMENT NUMBER:
                             112:2020
                  Plasmid composite for Escherichia coli and
  TITLE:
               thermophilic bacteria
                      Kawamata, Akiko; Fujita, Shozo; Asano, Takaharu;
  INVENTOR(S):
               Hataya, Takafumi
  PATENT ASSIGNEE(S): Fujitsu Ltd., Japan
                    Jpn. Kokai Tokkyo Koho, 3 pp.
  SOURCE:
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CODEN: JKXXAF

Japanese

DOCUMENT TYPE:

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

LANGUAGE:

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APPLICATION NO. DATE
  PATENT NO.
                   KIND DATE
  JP 01091784 A2 19890411 JP 1987-245852 19871001
AB ***Shuttle*** plasmids suitable for gene cloning in thermophilic
  bacteria at high temp. are prepd. from pBR322 of Escherichia coli and
  cryptic plasmid pTT8 of highly-thermophilic bacteria, e.g.
***Thermus***
  thermophilus. Plasmid pBTT1 and pBTT3 were prepd. from the two
  described above. The plasmids were maintained stably for >20
generations
  in T. thermophilus.
L6 ANSWER 17 OF 18 HCAPLUS COPYRIGHT 2002 ACS
ACCESSION NUMBER:
                            1989:34809 HCAPLUS
                             110:34809
DOCUMENT NUMBER:
                 A chloramphenicol-selectable plasmid and its
TITLE:
              construction for cloning in thermophilic bacteria
                      Yasuda, Hachiro; Fujita, Shozo; Asano, Takaharu;
INVENTOR(S):
              Kawamata, Akiko
PATENT ASSIGNEE(S): Fujitsu Ltd., Japan
                   Jpn. Kokai Tokkyo Koho, 3 pp.
SOURCE:
              CODEN: JKXXAF
DOCUMENT TYPE:
                          Patent
LANGUAGE:
                      Japanese
FAMILY ACC. NUM. COUNT: 1
PATENT INFORMATION:
   PATENT NO. KIND DATE
                                        APPLICATION NO. DATE
   JP 63148986 A2 19880621 JP 1986-296813 19861215
 AB The chloramphenicol acetyl transferase gene (CAT) is first successfully
   inserted into the plasmid pTT8 of ***Thermus*** thermophilus to
 obtain
   a chlorampheicol-selectable plasmid useful for cloning in thermophilic
   bacteria. A ***shuttle*** vector (no name given) for T. thermophilus
   and Escherichia coli was constructed by inserting in pTT8 a BamHI
   contg. the ampicillin-resistance gene of pBR322 and the CAT gene of
    plasmid pC194.
 L6 ANSWER 18 OF 18 HCAPLUS COPYRIGHT 2002 ACS
 ACCESSION NUMBER:
                             1983:66396 HCAPLUS
 DOCUMENT NUMBER:
                              98:66396
                   Construction of various host vector systems and the
 TITLE:
               variation of enzyme levels
                      Sakaguchi, K.
 AUTHOR(S):
                            Lab. Microbiol. Chem., Mitsubishi-Kasei Inst.
 CORPORATE SOURCE:
               Sci., Tokyo, Japan
                     Enzyme Eng. (1982), 6, 479-89
 SOURCE:
               CODEN: ENENDT; ISSN: 0094-8500
 DOCUMENT TYPE:
                           Journal
                        English
 LANGUAGE:
 AB Various published and unpublished expts. in the genetic engineering of
    several species are discussed, esp. the use of genetic engineering in the
    modification of enzyme expression. Expts. discussed include: the cloning
    of the Bacillus subtilis gene leu region on ***shuttle*** vector
    plasmids and subsequent expression in a Bacillus subtilis recE4 mutant,
    the increased activity of tryptophan synthetase [9014-52-2] encoded by
    Escherichia coli DNA cloned in Pseudomonas aeruginosa, the cloning and
    expression of the leuB and leuC genes which encode
  .beta.-isopropylmalate
    dehydrogenase [9030-97-1] and .alpha.-isopropylmate isomerase [50812-24-3], resp. of ***Thermus*** thermophilus in E. coli, the
     isolation of linear DNA plasmids from Streptomyces, protoplast fusion of
     Brevibacterium flavum, protoplast fusion of various yeast genera, the
     introduction of isolated yeast mitochondria into Saccharomyces cerevisiae
     protoplasts, and the introduction of whole cells of the N-fixing bacteria
     Azotobacter vinelandii and Anacystis nidulans into Saccharomyces
     cerevisiae protoplasts.
```

=> s 11 and 13 L7 3 L1 AND L3

=> dup rem 17

```
PROCESSING COMPLETED FOR L7
         3 DUP REM L7 (0 DUPLICATES REMOVED)
=> s 18 not 16
        2 L8 NOT L6
L9
=> d 19 ibib abs 1-2
L9 ANSWER I OF 2 HCAPLUS COPYRIGHT 2002 ACS
                           1997:748056 HCAPLUS
ACCESSION NUMBER:
DOCUMENT NUMBER:
                             128:137042
                  The Tsp45I restriction-modification system is
TITLE:
              plasmid-borne within its thermophilic host
                      Wayne, Jay; Holden, Megan; Xu, Shuang-yong
AUTHOR(S):
                            New England Biolabs Inc., 32 Tozer Road,
CORPORATE SOURCE:
Beverly, MA
                    Gene (1997), 202(1/2), 83-88
SOURCE:
               CODEN: GENED6; ISSN: 0378-1119
PUBLISHER:
                      Elsevier Science B.V.
DOCUMENT TYPE:
                          Journal
                       English
LANGUAGE:
                      species YS45 harbors two small cryptic plasmids of
      ***Thermus***
   (pTsp45s) and approx. 12 kb (pTsp45l). Plasmid pTsp45s has been
entirely
   sequenced, revealing three significant ORFs. In addn. to a previously
   reported thermophilic plasmid-encoded replication protein (Rep), pTsp45s
   contains two genes for the Tsp45I methyltransferase (M.Tsp45I) and
    restriction endonuclease (Tsp45I). These two converging genes (tsp45IM
    and tsp45IR) overlap by 4 bp at their stop codons within an Xbal site.
    M.Tsp45I (413 aa, 47.0 kDa, recognizing 5'-GTSAC-3') is highly
 homologous
    to other m6A-methyltransferases, esp. M.Ecal (recognizing
 5'-GGTNACC-3').
    Tsp45I (332 aa, 37.4 kDa, cleaving 5'-.dwnarw.GTSAC-3') is not
 homologous
    to M.Tsp45I, or to other restriction endonucleases. Recombinant Tsp45I
    stably produced in E. coli, and cleaves DNA at 65.degree.C with the same
    specificity as the native enzyme. Therefore, the thermophilic Tsp45I
    restriction-modification system is plasmid-borne within its native host.
 L9 ANSWER 2 OF 2 HCAPLUS COPYRIGHT 2002 ACS
                             1997:504779 HCAPLUS
 ACCESSION NUMBER:
 DOCUMENT NUMBER:
                              127:230135
                   Identification of a thermophilic plasmid origin and
 TITLE:
               its cloning within a new ***Thermus*** -E. coli
                       Wayne, Jay; Xu, Shuang-yong
 AUTHOR(S):
                             New England Biolabs, 32 Tozer Road,
  CORPORATE SOURCE:
 Beverly, MA,
                01915, USA
                     Gene (1997), 195(2), 321-328
  SOURCE:
                CODEN: GENED6; ISSN: 0378-1119
  PUBLISHER:
                       Elsevier
  DOCUMENT TYPE:
                            Journal
  LANGUAGE:
                        English
  AB A pUC19-based vector has been generated for selecting functional
    thermophilic origins (oris) of ***Thermus*** ssp. Once combined with
    thermophilic DNA, the vector can be amplified in ampicillin resistant
     (ApR) E. coli, prior to transformation and kanamycin (Km) selection in
      ***Thermus*** thermophilus. The KmR ***Thermus***
    replicate any newly-formed shuttle vectors via introduced thermophilic
    oris. Using this 'ori-selecting' vector, three novel thermophilic oris were cloned from randomly digested ***Thermus*** cryptic plasmid
  DNA.
     These shuttle vectors are useful for genetic analyses, as well as protein
     engineering within thermophiles. The smallest ori-contg. sequence of
     4.2kb has been subcloned, sequenced, and further refined to 2.3kb. A
     significant ORF of 341 amino acids (aa), with a ***Thermus***
     and RBS, is found within the thermophilic ori. Deleting part of this ORF
     abolishes the shuttle vector's ability to replicate in T. thermophilus.
     Therefore, we postulate that this ORF encodes a replication protein (Rep)
```

necessary for thermophilic plasmid replication. The thermophilic ori also

contains two sequences which resemble DnaA boxes.